

iGEM 2016 – Microbiology – BMB – SDU

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| Project type: Silk MaSp2 BC | Creation date: 2016.09.21 |
| Project title: Cloning composite part into iGEM standard plasmid psB1C3. | Written by: Rune Øbo |
| Sub project: Insertion of K1763003 (MaSp2 AB) into pSB1C3 plasmid | Performed by: Rune Øbo, Nete Sloth Bækgaard, Mathilde Nygaard, Rikke Friis Bentzon & Viktor Swaglord Mebus. |

1. SOPs in use.

SOP number: SOP0007_v01 LA plates with antibiotic

SOP number: SOP0022_v01 Competent cell - freeze-stock

SOP number: SOP0023_v01 Ca⁺⁺ transformation

SOP number: SOP0009_v01 TSB transformation

Plasmid purification kit

SOP number: SOP0001_v01 ON culture of *E.coli*

SOP number: SOP0004_v01 Bacterial freeze stock

SOP number: SOP0017_v01 Fast digest

SOP number: SOP0015_v01 Ligation

Gel purification kit

SOP number: SOP0021_v01 Colony PCR with MyTaq

2. Purpose.

To insert basic part: K1763003 (MaSp2 BC), into iGEM standard plasmid; pSB1C3.

3. Overview.

| Day | SOPs | Experiments |
|-----|----------------------|---------------------------------|
| 1 | SOP0023_v01 | Ca ⁺⁺ transformation |
| 2 | SOP0001_v01 | ON culture of <i>E.coli</i> |
| 3 | Miniprep kit | Plasmid purification |
| 3 | SOP0001_v01 | ON culture of <i>E.coli</i> |
| 4 | SOP0004_v01 | Bacterial freeze stock |
| 5 | SOP0017_v01 | Fast digest |
| 5 | Gel purification kit | Gel purification |
| 6 | SOP0015_v01 | Ligation |
| 6 | SOP0009_v01 | TSB Transformation |
| 7 | SOP0021_v01 | Colony PCR with MyTaq |
| 7 | SOP0001_v01 | ON culture of <i>E.coli</i> |
| 8 | Miniprep kit | Plasmid purification |
| 8 | SOP0017_v01 | Fast Digest with EcoRI + PstI |
| 9 | SOP0004_v01 | Bacterial freeze stock of ON. |

4. Materials required.

Materials in use

| Name | Components (Concentrations) | Manufacturer / Cat. # | Room | Safety considerations |
|---------------------------------|--|---------------------------------|----------------------------------|--------------------------|
| Appropriate medium ex. LB | 1% Tryptone 1% NaCl 0.5% Yeast extract | Oxoid Sigma-Aldrich Merck | Media lab or V18-40 5-0 | |
| Glycerol | 50 % | AppliChem | Anne Mette, RT | |
| LB | | Anne-Mette | | |

| | | | | |
|--|---|---|-------------------------------|-------------------|
| LA | 1% Tryptone 1% NaCl 0.5% Yeast extract 1.5% agar | Oxoid Sigma-Aldrich Merck Difco agar from BD | Anne-Mette Or V18-405-0 | |
| Water | Demineralised milli-Q autoclaved water | Milli-Q water purification system (Millipore) | RT | Water |
| MyTaq™HS Red Mix | http://www.bioline.com/documents/product_inserts/MyTaq%E2%84%A2%20HS%20Red%20Mix.pdf#zoom=130 | Bioline | V18-405a-2 | |
| Reverse primer | Made specific to the template | Sigma-Aldrich | | |
| Forward primer | Made specific to the template | Sigma-Aldrich | | |
| Ligasebuffer | | Agilent Technologies | Freezer at 1. Floor | |
| Ligase | | | Freezer 1. Floor | Ligase |
| FastDigest enzyme | | Agilent Technologies | Freezer at 1. Floor | EcoRI PstI |
| Fast digest green / 10 x FastDigest Buffer | | Agilent Technologies | Freezer at 1. | |
| CaCl ₂ | 0.1M | | Chem room | |
| MgCl ₂ | 0.1M | | Chem room | MgCl ₂ |
| liquid nitrogen | liquid nitrogen | liquid nitrogen | liquid nitrogen | |
| 6x DNA Loading Dye | | GeneRuler | fridge floor 1 | |
| Fort. LB | | The new Anne-Mette | Autoclave room | |
| Magnesium chloride (MgCl ₂) 1M | 1M | The New Anne-Mette | Autoclave | |

| | | |
|---------------------------------|---------------|---------------------|
| Polyethylene glycol (PEG) 3.350 | Sigma Aldrich | Micro Chemical room |
| Dimethylsulfoxid (DMSO) | Sigma Aldrich | Micro Chemical room |
| | | |

5. Other

As competent cells, LB and LA media was used by all parts of our project and not just this protocol the dates for use of these SOPs are not added. this comment deal with SOP number: SOP0007_v01 and SOP0022_v01

Gel Electrophoresis is set at 75 V for 30-45 minutes, dependent on the gel percent.

6. Experiment history.

| Date (YY.MM.DD) | SOPs | Alterations to SOPs and remarks to experiments |
|-----------------|----------------------------|--|
| 16.06.27 | | Resuspend the received 200ng gblock from IDT with dH2O. Thereafter take 5µl of diluted gblock and add 95µl dH2O making a 20X dilution. (K1763003) Name: SG7 |
| 16.07.22 | SOP0017_v01 Fast Digest | Here we used the Bacto-groups plasmid with inserted bactericidine, lysostaphin. Fast Digest with EcoRI for 30min and PstI for 2h. Gel electrophoresis. |
| 16.07.22 | Gel Purification kit. | Ran the gel electrophoresis for 40min 75V. Cut out and purified the backbone psB1C3. Name SG58. |
| 16.07.22 | SOP0010_v01 Phusion PCR | Phusion PCR of SG7. Program used for PCR: |

| Temp (°C) | Time (min) |
|-----------|------------|
| 95°C | 3:00 |
| 95°C | 0:25 |

| | |
|---------|------------|
| 58°C | 0:25 |
| 72°C | 0:20 |
| Repeats | x34 |
| 72°C | 1:00 |
| 20°C | Infinite . |

Name: SY12

16.07.22 SOP0017_v01 Fast Digest Fast Digest Fast Digest of SY12 with PstI for 2h at 37°C and EcoRI for 0.5h at 37°C. After digestion the enzymes were denaturated at 80°C in 10min.
Name: SG48

16.07.22 SOP0015_v01 Ligation Ligation of SG58 (digested backbone) and SG48 (digested gblock). Ligation took place at 16°C for 16h.
Name: SB73

| | | |
|---------------|-------|-------|
| Ratio | 1:2 | 1:5 |
| Ligase Buffer | 2µl | 2µl |
| Ligase | 1µl | 1µl |
| SG58 | 1µl | 1µl |
| SG48 | 2µl | 5µl |
| H2O | →20µl | →20µl |

16.07.23 Enzyme inactivation Denaturation of enzyme ligase with 65°C for 10min.

16.07.23 SOP0009_v01 TSB transformation 16.07.22 SOP0001_v01 ON culture of *E. coli*. Top 10. OD about 0.3-0.5nm. Thereafter a regular TSB transformation with SB73.
Plating after TSB transformation.

16.07.24 SOP0021_v01 Colony PCR with MyTaq Taking 3 colonies from the plate to increase the odds of one colony to be correct. Primers VR and VF2. Using the same pipet tip, the colonies were also set to an ON.
Half of the colony PCR product were added EcoRI and PstI to see if they were able to recognize. Gene around 119bp.

| | |
|------|------------|
| Temp | Time (min) |
|------|------------|

| | | | |
|----------|---|---|----------|
| | | 95°C | 3:00 |
| | | 95°C | 0:25 |
| | | 58°C | 0:25 |
| | | 72°C | 0:30 |
| | | Repeats | 34X |
| | | 72°C | 1:00 |
| | | 20°C | Infinite |
| 16.07.24 | Gel electrophoresis | 2% agarose gel, 75V for 30min. | |
| 16.07.25 | SOP0004_v01 Bacterial Freeze stock | 900µl of the previous ON culture were added into 125µl glycerol and put in the -80°C freezer. Name: #41 | |
| 16.07.25 | Miniprep Kit (plasmid purification kit) | Plasmid purification on the same ON culture as freeze stock. Name: SR39. | |
| 16.07.26 | Sequencing | Sent SR39 to sequencing. | |

7. Sample specification.

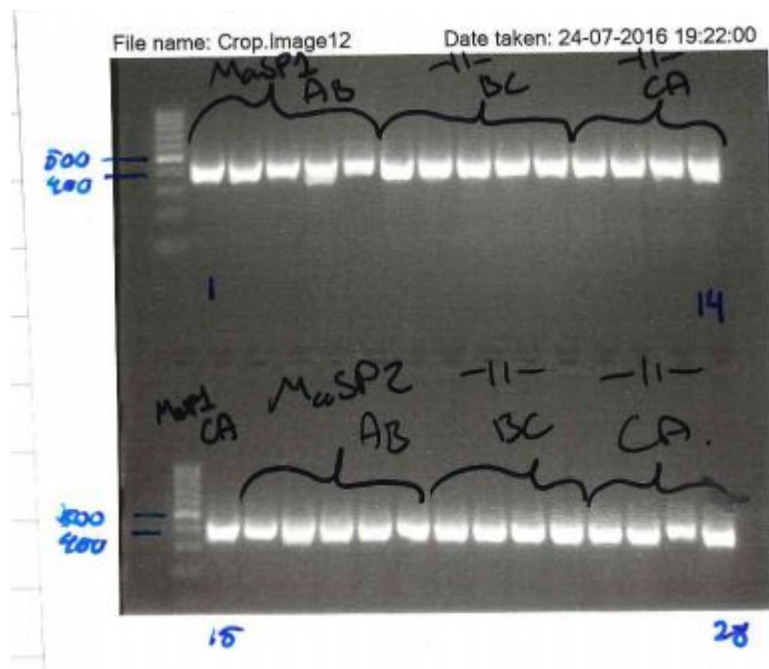
| Sample name | Sample content | From | Used for / Saved where |
|-------------|----------------------------|------|--|
| SG7 | MaSp2 BC g block: K1763003 | IDT | Diluted with dH2O / saved in coolbox |
| SY12 | Phusion PCR product of SG7 | IDT | Used to cut with restriction enzymes EcoRI and PstI. |
| SG58 | purified psB1C3 | iGEM | Vector cut with pstI and EcoR1 / cool box |
| SG48 | Fast digest of SY12 | | Cut with PstI and EcoR1 / Saved in coolbox |

| | | |
|------|----------------------------|--|
| SB73 | pSB1C3 ligated with SG48 | vector and basic part ligated and transferred to <i>E.coli</i> . /Saved in coolbox |
| SR39 | pSB1C3:k1763003 (MaSp2 BC) | purified and transferred SB73/ freeze stock in <i>E.coli</i> (#41) |

8. Remarks on setup.

9. Results and conclusions.

The first gel photo below shows the colony PCR results. The second gel photo shows cutting of SB75 with Pst1 at 2 hours with 37 °C and EcoR1 at 30 minutes, and after that called SR40. Every second well had added restriction enzymes. The composite part is 119bp. It is possible from the Gel photo to assume that the DNA is cut correctly. VR and VF2 primers adds a total of 300bp, so a total of around 419bp. From this gel the ligation and transformation of SB75, band with around 119bp after digestion, were concluded successfully and sent to sequencing.



Picture of colony PCR of different colonies on the same plate, but each gene had its own plate. All the colony PCR shows a band near the 500makr which was desired.



This gel photo shows the cutting of either PstI and also another cut with PstI and EcoRI. Every second well had added two restriction enzymes. The markings in between the 200 mark and 100mark of base pairs indicates our correct gene. The first 6 wells are MaSp1 and the last 6 are the genes for MaSp2. Ladders were added on both sides, blue and green.

10. Appendixes